CARDIOVASCULAR MEDICINE

Markers of inflammation and multiple complex stenoses (pancoronary plaque vulnerability) in patients with non-ST segment elevation acute coronary syndromes

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Objective: To assess the relation between markers of inflammation and the presence of multiple vulnerable plaques in patients with non-ST segment elevation acute coronary syndromes.

Design: Prospective cohort study of 55 patients with non-ST segment elevation acute coronary syndromes and angiographically documented coronary disease. Blood samples were obtained at study entry for the assessment of high sensitivity C reactive protein (CRP), neopterin, and neutrophil count. Coronary stenoses were assessed by quantitative computerised angiography and classified as "complex" (irregular borders, ulceration, or filling defects) or "smooth" (absence of complex features). Extent of disease was also assessed by a validated angiographic score.

Results: Neutrophil count (r = 0.36, p = 0.007), CRP concentration (r = 0.33, p = 0.02), and neopterin concentration (r = 0.45, p < 0.001) correlated with the number of complex stenoses. Patients with multiple (three or more) complex stenoses, but not patients with multiple smooth lesions, had a higher neutrophil count ($5.9 (1.4) \times 10^9/l \times 4.8 (1.4) \times 10^9/l$, p = 0.02), CRP concentration (log transformed) ($1.08 (0.63) \times 0.6 (0.6)$, p = 0.03), and neopterin concentration (log transformed) ($0.94 (0.18) \times 0.79 (0.15)$, p = 0.002). Multiple regression analysis showed that neopterin concentration (B = 4.8, 9.5% confidence interval (CI) 1.9 to 7.7, p = 0.002) and extent of coronary artery disease (B = 0.6, 9.5% CI 0.03 to 1.2, p = 0.04) were independently associated with the number of complex stenoses.

Conclusions: Acute inflammatory markers such as high neutrophil count, CRP concentration, and neopterin concentration correlate with the presence of multiple angiographically complex coronary stenoses. Neopterin concentration was a stronger predictor of multiple complex plaques than were neutrophil count and CRP concentration. These findings suggest that a relation exists between inflammation and pancoronary plaque vulnerability.

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cute coronary syndromes (ACS) are usually associated with "vulnerable" atheromatous plaques, which are prone to fissuring or to developing endothelial erosion.1 2 At angiography, disrupted or ulcerated plaques often appear as "complex" stenoses with rough contours or "filling defects" suggestive of intracoronary thrombosis.³⁻⁶ The presence of multiple complex coronary plaques is known to be associated with adverse prognosis in patients with coronary artery disease.7 Inflammation appears to play an important part in plaque vulnerability and disruption. White blood cell activation and the subsequent release of proinflammatory cytokines can promote rupture or erosion of atheromatous coronary plaques through a number of mechanisms.12 Inflammatory markers such as C reactive protein (CRP), neutrophil count, and neopterin are increased in patients with ACS and are markers of coronary artery disease activity.8 9 Moreover, previous reports from our group showed that neopterin concentration correlates with the presence of complex stenoses in patients with unstable angina.10

Previous work by Buffon and colleagues¹¹ and Rioufol and associates¹² suggests that ACS is a pancoronary condition associated with multiple plaque vulnerability. We therefore sought to study the relation between inflammatory markers and the presence of multiple complex (vulnerable) plaques—that is, those most at risk of recurrent events in patients presenting with ACS.

METHODS

Patients

We studied 55 of 106 consecutive patients who were admitted to the coronary care unit of St George's Hospital in London between September 1996 and June 1997 with non-ST segment elevation ACS and whose condition stabilised rapidly with medical treatment. All patients in the study had angiographically documented coronary artery disease and had no evidence of local or systemic inflammatory conditions. Excluded were 40 patients without angiography, seven patients with systemic inflammatory conditions, and four with creatinine concentration ≥ 120 mmol/l. The latter were excluded because renal function is a major determinant of blood neopterin concentration. Cardiac troponin was measured for only 32 patients, as this marker was not routinely measured in our institution at the time of the study. The patients' clinical management and the decision to proceed to cardiac catheterisation were left to the discretion of the managing cardiologists, who were unaware of results regarding inflammatory markers. Some patients in the present study had been enrolled in a previous study by our group.10 All patients had primary unstable angina, corresponding to subclass "b" of Braunwald's classification. 13 Forty seven patients were in Braunwald class IIIb and eight were in class IIb. Cardiac troponin T was abnormal in 10 of the 32 patients tested. All patients had transient ECG changes that were diagnostic of myocardial ischaemia and

received treatment with aspirin (75–325 mg), heparin, intravenous glyceryl trinitrate, and β blockers. All patients gave written informed consent before study entry and the study was approved by the local research ethics committee.

Blood sampling, neutrophil count, CRP, and neopterin measurements $\,$

Blood samples from every patient were drawn at admission to the coronary care unit. Neutrophils were counted by an automated blood cell counter. Blood for CRP measurement was drawn and centrifuged immediately and the serum aliquoted and stored at −80°C until analysis. No specimen inadvertently thawed during storage. CRP was measured on the COBAS Integra (Roche Diagnostics Limited, Lewes, East Sussex, UK) with the CRP latex assay in both the high sensitivity application (analytical range 0.2-12 mg/l) and the normal application (analytical range 2-160 mg/l). The analytical precision of the high sensitivity CRP latex assay was 7.6% at a concentration of 1.02 mg/l, 3.3% at 1.79 mg/l, and 1.3% at 4.36 mg/l. Samples outside the analytical range of the high sensitivity CRP latex assay were analysed by the CRP latex assay in the normal application. The analytical precision of the normal CRP latex assay was 2.4% at a concentration of 29.5 mg/l and 1.3% at 113 mg/l. Serum neopterin concentration was measured with a commercially available immunoassay (ELISA kit, IBL, Hamburg, Germany). The within coefficient variability in the 7.7 nmol/l range was less than 3% and was below 4% in the 20 nmol/l

Angiographic analyses

Coronary angiography was carried out according to the Judkins technique. Images of the coronary tree of all patients were obtained in routine, standardised projections with the digital quantitative Philips Integris 3000 system (Philips, the Netherlands). Two experienced cardiologists who had no knowledge of the patients' clinical characteristics and biochemical results visually reviewed all angiographic images to assess the extent of coronary artery disease and to characterise the morphology of all coronary artery stenoses with \geqslant 30% diameter reduction.

Angiographic scoring system

To assess coronary artery disease extent and severity, angiograms were scored according to the number of major coronary arteries with \geq 75% reduction in lumen diameter (vessel score). Interobserver variability (the standard deviation of the mean unsigned difference between paired estimates) for the vessel score in this study was 4.9%.

Angiographic coronary stenosis morphology

Lesion morphology was assessed in all stenoses with > 30% diameter reduction as reported in previous studies by our group and others.14-19 Briefly, stenoses were subdivided into complex or smooth. Complex lesions were defined by the following features: (a) irregular morphology, scalloped borders, or both; (b) overhanging or abrupt edges perpendicular to the vessel wall; (c) ulceration (that is, outpouchings within the stenosis); and (d) the presence of filling defects consistent with intracoronary thrombus. Of interest, 88% of coronary stenoses defined as complex corresponded to type C in the American College of Cardiology/American Heart Association classification.²⁰ Coronary stenoses with no complex features were classified as smooth lesions. When discrepancies arose regarding the morphological appearance of a lesion, a third experienced observer examined the stenosis and the lesion was classified by consensus. The reproducibility of the findings was determined by repeat analysis more than three months later by two experienced observers who had no knowledge of the results of stenosis classification obtained at the first reading. Interobserver agreement regarding qualitative morphological analyses was 97%

Statistical analysis

Normality was analysed with the Kolmogorov-Smirnov test. CRP and neopterin serum concentrations had a non-normal distribution and were therefore log transformed before analysis. Results for normally distributed continuous variables are expressed as mean (SD) and continuous variables with non-normal distribution are presented as median value (interquartile interval). Comparisons of continuous variables were analysed by the unpaired t test. The Spearman two way test was used to assess the relation between two quantitative variables with non-normal distribution. The Pearson two way test was used to assess the relation between two quantitative variables with normal distributions. We assessed independent predictors of complex stenoses with multiple regression analysis. In the multiple regression analysis the dependent variable was the number of complex lesions; the independent variables were those that, on univariate analysis, correlated significantly with the number of complex lesions and variables that showed a trend (p < 0.20) towards an association. Differences were considered to be significant if the null hypothesis could be rejected with > 95% confidence. The SPSS 11.5 statistical software package (SPSS Inc, Chicago, Illinois, USA) was used for all calculations.

RESULTS

Table 1 shows the baseline clinical and biochemical characteristics and table 2 the angiographic results of the 55 patients in the study. Complex lesions were found in 44 (80%) patients, of whom 15 had multiple (three or more) complex lesions in one or more coronary arteries. Smooth

Table 1 Baseline clinical and biochemical characteristics of 55 patients with non-ST segment elevation acute coronary syndromes and angiographically documented coronary artery disease

	Mean (SD) or n (%)
Age (years)	62.5 (10.5)
Male sex	41 (74.5%)
Systolic blood pressure (mm Hg)	138.9 (19. <i>7</i>)
Diastolic blood pressure (mm Hg)	77.1 (14.3)
Body mass index (kg/m²)	26.8 (4.1)
Previous PTCA	14 (25.5%)
Previous myocardial infarction	31 (56.4%)
Previous unstable angina	28 (50.9%)
Previous stroke	3 (5.5%)
Cardiovascular risk factors	
Diabetes mellitus	7 (12.7%)
Hyperlipidaemia*	27 (49.1%)
Hypertension	29 (52.7%)
Smoking	14 (25.5%)
Biochemistry	
Cholesterol (mmol/l)	5.4 (1)
HDL cholesterol (mmol/l)	1 (0.3)
LDL cholesterol (mmol/l)	3.4 (0.8)
Triglycerides (mmol/l)	1.7 (1)
Treatment	
Aspirin	40 (72.7%)
β Blockers	24 (43.6%)
Nitrates	27 (49.1%)
Calcium channel antagonists	30 (54.5%)
ACE inhibitors	10 (18.2%)
HMG-CoA reductase inhibitors	13 (23.6%)

^{*}Total cholesterol > 5.4 mmol/l.

ACE, angiotensin converting enzyme; HDL, high density lipoprotein; HMG-CoA, hydroxymethyl glutaryl coenzyme A; LDL, low density lipoprotein; PTCA, percutaneous transluminal coronary angioplasty.

Table 2 Inflammatory markers and angiographic variables in 55 patients with acute coronary syndromes

Variable	Mean (SD) or median (IQR)
Inflammatory markers	
White blood cell count (× 10 ⁹ /l)	8 (1.7)
Neutrophil count (× 10 ⁹ /l)	5.1 (1.5)
Log ₁₀ neopterin	0.81 (0.03)
Log ₁₀ CRP	0.7 (0.64)
Angiography	, ,
Vessel score	2.0 (1-3)
Number of smooth stenosis	3.5 (1.8)
Complex stenosis (units)	1.7 (1.5)
Ejection fraction (%)	59.1 (14.9)

lesions were observed in 53 (96.4%) patients, 35 of whom had multiple (three or more) smooth lesions.

Neutrophil count, CRP concentration, and neopterin concentration correlated significantly with the number of complex stenoses but not with the number of smooth coronary lesions (table 3). Of the three markers assessed, neopterin was most strongly correlated with the number of complex stenoses. As neopterin concentration is known to be influenced by age, neopterin concentrations were adjusted by age and reanalysed.

Patients with multiple complex stenoses had higher neutrophil count, CRP concentration, and neopterin concentration than patients with fewer than three complex lesions. Concentrations of the three inflammatory markers where similar in patients with multiple smooth lesions and patients with fewer than three smooth stenoses (fig 1).

Univariate analysis showed that age, low density lipoprotein cholesterol concentration, and vessel score were significantly correlated with the number of complex lesions in our patients (table 4). Backward stepwise multiple regression analysis was performed with, firstly, all three inflammatory markers; secondly, variables significantly related to complex stenosis on univariate analysis; and thirdly, variables showing a trend (p < 0.20) towards an association. Neopterin concentration and vessel score were the only two variables that remained independent predictors of the number of complex lesions in this model (table 5). Multivariate analysis, without the inclusion of neopterin in the model, showed that neutrophil count and vessel score, but not CRP, were independent predictors of the number of complex lesions (data not shown).

Table 3 Univariate correlation between neutrophil count, CRP concentration, and neopterin concentration (both log transformed), and angiographic data

	r	p Value
Neutrophil count		
Vessel score*	0.10	0.43
Number of complex stenoses	0.36	0.007
Number of smooth stenoses	0.03	0.81
Ejection fraction (%)	0.16	0.29
Log 10 CRP		
Vessel score*	0.26	0.07
Number of complex stenoses	0.33	0.02
Number of smooth stenoses	0.12	0.40
Ejection fraction (%)	-0.003	0.98
Log10 neopterin†		
Vessel score*	0.09	0.51
Number of complex stenoses	0.45	< 0.001
Number of smooth stenoses	0.004	0.97
Ejection fraction (%)	0.13	0.4

Two way Pearson or *Spearman correlation test; †adjusted by age.

DISCUSSION

The main finding of the present study is that the presence of multiple vulnerable plaques is associated with increased concentrations of inflammatory markers. These findings suggest a link between inflammation and pancoronary plaque vulnerability. In our study, inflammatory markers such as neutrophil count, CRP, and neopterin correlated significantly with multiple plaque complexity. Neopterin was the strongest predictor of multiple plaque complexity. Thus, inflammatory markers as measured in our study identify a vulnerable phenotype among patients with non-ST segment elevation ACS.

Inflammatory markers and angiographic complexity

A previous study from our group showed that neopterin concentration is a predictor of the presence of complex stenoses in patients with unstable angina.10 A recent study from Zairis and colleagues²¹ reported that CRP concentration correlated with stenosis complexity in patients with unstable angina. They studied 228 patients with primary unstable angina and found that CRP concentration was independently associated with the presence of either multiple stenosis or angiographically apparent thrombus containing complex lesions. Katritsis and colleagues²² linked CRP concentration to various angiographic characteristics in 103 patients undergoing cardiac catheterisation for suspected coronary artery disease. They found higher CRP concentration among patients with macroscopic thrombus without total occlusion of the vessels than among patients without thrombus and total vessel occlusion. Sabatine and colleagues23 correlated angiographically apparent thrombus with increasing white blood cell count in patients with unstable angina. Unfortunately, they did not assess the relation between neutrophil count and plaque complexity. The present study is the first to assess specifically the relation between three inflammatory markers and the presence of multiple angiographically complex lesions in patients with non-ST segment elevation ACS. Our study lends support to the hypothesis that inflammation may be responsible for a pancoronary process leading to the development of multiple complex lesions in individual patients. We showed here that neutrophil count, CRP, and neopterin were predictors of multiple angiographically complex lesions suggestive of plaque disruption, with neopterin being the strongest predictor after adjusting for confounding factors. Of importance, no differences were found in inflammatory marker concentrations between patients with and those without multiple smooth lesions. This finding further strengthens the argument that inflammation influences the presence of a vulnerable phenotype and this in turn appears to be a pancoronary process.

Inflammation, macrophages, and neopterin

Findings in this study confirm and expand previous observations from our group regarding neopterin as a predictor of multiple complex lesions in patients with unstable angina.10 Several studies have previously shown that macrophages play a key part in destabilising atheromatous plaques. 1 2 24 Neopterin is a pteridine derivative that is secreted by macrophages after stimulation by interferon γ . 25 It has been suggested that complex lesions seen on angiography are vulnerable plaques prone to disruption or actually disrupted plaques. 14 26 27 Plaque vulnerability has been shown to be a function of the increased local number of inflammatory cells within plaques, particularly macrophages and activated lymphocytes.28 Activated lymphocytes in atherosclerotic plaques produce interferon γ , a molecule that reduces collagen synthesis within atheromatous plaques and activates macrophages. Activated macrophages synthesise and release matrix metalloproteinases, which contribute to

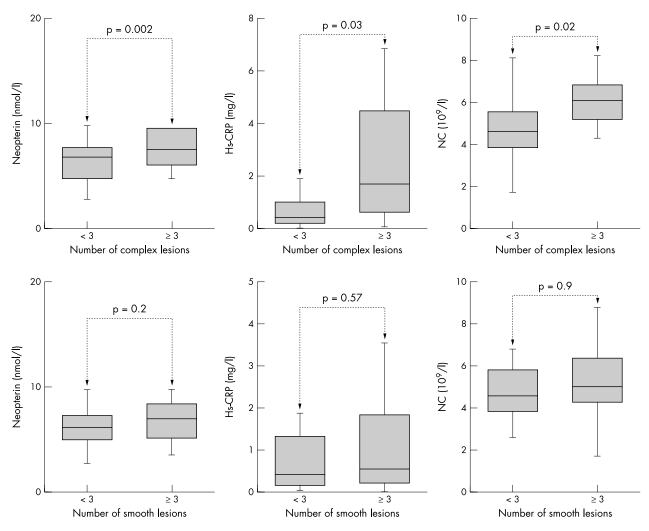


Figure 1 Box plots showing differences in neopterin concentration, high sensitivity C reactive protein (Hs-CRP) concentration, and neutrophil count (NC) between patients with multiple complex lesions (≥ 3) and those without, and between patients with multiple smooth lesions (≥ 3) and those without.

the degradation of collagen in the fibrous cap.²⁹ These processes result in an imbalance between synthesis and degradation of matrix components in areas of active inflammation within atherosclerotic plaques ("vulnerable sites"), which favours plaque rupture.^{30 31} Macrophages activated by interferon γ synthesise neopterin,²⁵ which, in

turn, can interfere with intracellular signalling pathways known to be influenced by oxidative stress. 32 33 Neopterin activates the redox sensitive transcription nuclear factor κB , 34 which upregulates proinflammatory genes such as interleukin 6 and tumour necrosis factor α . 35 The overall result is an increase of the inflammatory tone within the vascular wall. 36

Table 4 Univariate correlation (*r*) between baseline variables and complex coronary artery stenosis

	r	p Value
Age (years)	0.42	0.001
Body mass index (kg/m²)	0.04	0.78
Systolic blood pressure (mm Hg)	0.18	0.18
Diastolic blood pressure (mm Hg)	-0.02	0.88
Biochemistry		
Cholesterol (mmol/l)	0.10	0.44
HDL cholesterol (mmol/l)	-0.16	0.26
LDL cholesterol (mmol/l)	0.34	0.02
Triglycerides (mmol/l)	-0.005	0.97
White blood cell count (× 10 ⁹ /l)	0.10	0.45
Angiography		
Vessel score*	0.60	< 0.001
Ejection fraction (%)	-0.22	0.13
Number of smooth lesions	-0.07	0.62

Widespread inflammation in unstable angina

Our results confirm and expand previous observations by others¹¹ ¹² ³⁷ that ACS is a pancoronary condition related to

 Table 5
 Multiple regression analysis with univariate predictive variables for number of complex lesions

Variable	B (95% CI)	t	p Value
Neopterin (log transformed)	4.8 (1.9 to 7.7)		0.002
Vessel score	0.6 (0.03 to 1.2)		0.04

Neutrophil count (p=0.1), CRP concentration (p=0.42), LDL cholesterol (p=0.49), left ventricular ejection fraction (p=0.32), age (p=0.71), systolic blood pressure (p=0.36), and treatment with β blockers (p=0.79) and lipid lowering drugs (p=0.95) were not independent predictors of complex lesions.

B, increment of the number of complex lesions with every unit of the independent variable; CI, confidence interval; t, statistic of each variable in the equation.

the presence of multiple vulnerable plaques. Inflammation and pancoronary plaque vulnerability appear to be directly related and the results of our study give further support to the importance of systemic and local inflammation in multiple atheromatous plaque disruption. Indeed, neutrophil count, neopterin concentration, and CRP concentration were higher in patients with multiple complex lesions than in those with fewer than three complex stenoses. Our findings provide a pathophysiological basis to recent reports by Goldstein et al,7 Rioufol et al,12 and Asakura et al,37 who observed multiple plaque fissuring in patients with ACS. Goldstein et al7 showed with coronary angiography the existence of additional unstable lesions other than the culprit lesions in 21% of patients with myocardial infarction. Using intravascular ultrasound, Rioufol et al12 found culprit lesion fissuring in over 37% of patients presenting with a first ACS and troponin I increase. In 79% of these patients at least one plaque rupture was found in stenoses other than the culprit lesion. Thus, ACS seems to be associated with pancoronary instability. Asakura et al³⁷ carried out angioscopic studies in patients undergoing catheterisation one month after the onset of myocardial infarction and observed that all three major coronary arteries were widely diseased and had multiple vulnerable plaques.

Our findings that neutrophil count, neopterin concentration, and CRP concentration are raised in patients with multiple complex stenoses suggest that systemic inflammation may be responsible for this widespread involvement of the coronary tree in patients with ACS. Moreover, our data are in agreement with findings of Buffon *et al*, " who found generalised activation of neutrophils across the coronary vascular bed in patients with unstable angina, regardless of the location of the culprit stenoses. In agreement with these findings, Naruko *et al*; " recently found in coronary artery segments from patients with myocardial infarction the occurrence of neutrophil infiltration in ruptured and eroded plaques as well as a higher number of activated neutrophils in patients with ruptured plaques than in patients with eroded plaques.

Limitations of the study

The present study design did not allow us to determine the mechanisms responsible for higher neutrophil count, CRP concentration, and neopterin concentration in patients with multiple complex lesions compared with those without. However, in the light of results obtained by Buffon et al111 and Naruko et al,38 it is conceivable that local and systemic inflammation linked to vulnerable plaques could have resulted in raised concentrations of inflammatory markers in our patients. Only a relatively small proportion of patients received statins, probably reflecting standard clinical practice at the time of patient recruitment. Another limitation is that coronary angiography is a "lumenogram" and therefore provides no information regarding phenomena that occur within the vessel wall. Coronary angiography thus has limitations compared with angioscopy and intravascular ultrasound. Moreover, Maehara et al39 have recently reported a strong correlation between the angiographic features of coronary plaque rupture and ultrasound findings in patients with unstable and stable coronary syndromes. Waxman et al40 found that angiographically complex atherosclerotic lesions are highly specific markers of plaque instability. Although correlations between inflammatory markers and stenosis morphology are significant in the present study, the clinical significance of this finding requires further investigation.

Conclusions

Our results indicate a relation between markers of inflammation and pancoronary plaque vulnerability in patients with

ACS. Acute inflammatory markers such as neutrophil count, CRP concentration, and neopterin concentration are higher among patients with multiple angiographically complex plaques than among those without and may help to identify a phenotype with increased plaque vulnerability.

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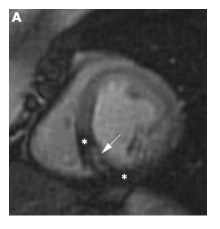
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IMAGES IN CARDIOLOGY.....

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Impending septal rupture in myocardial infarction detected by cardiac magnetic resonance imaging

67 year old woman was admitted one week after an ambulatory inferior myocardial infarction. Cardiac magnetic resonance imaging (MRI) (1.5 T, Sonata, Maestro class; Siemens, Erlangen, Germany) was performed the next day to assess myocardial viability. Contrast enhanced dynamic myocardial perfusion imaging (TrueFISP-sequence) at rest demonstrated contrast flow from the cavity of the left ventricle into the myocardium near the basal septum (arrows). The contrast agent penetrated about three quarters of the myocardial thickness, thus showing impending myocardial rupture (panel A). Using late enhancement technique (TurboFLASHsequence) 10 minutes after application of contrast agent, an extended transmural inferior infarction could be seen in the long axis view (panel B). Impending myocardial rupture is nontransmural and does not reach the subepicardial fatty tissue (†). Inside the myocardial infarction a zone with hypointense signal represented microvascular obstruction (*). Four hours after the MRI examination the patient





developed cardiogenic shock with a loud systolic murmur. Left ventricular angiography in a left anterior oblique (LAO) 60° position demonstrated an inferiorly located ventricular septal defect (panel C).

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